Marked-up Version Substitute Specification

DESCRIPTION TITLE OF THE INVENTION

RUBBER COMPOSITION

Rubber Composition

CROSS-REFERENCE TO RELATED APPLICATION

This application is a section 371 of International Application No. PCT/UF2005/023379, filed December 20, 2005, which was published in the Japanese language on May 18, 2006, under International Publication No. WO 2006/052035 Al and the disclosure of which is incorporated herein by reference.

Technical Field

[0001]

The present invention relates to a silica compounded rubber composition suitable for tire cap treads, which has a small die swell and excellent extrusion processability while keeping excellent wet skid performance, exothermic characteristic and abrasion resistance as required for the performance of tires such as safety and economy.

Also, the invention relates to a rubber composition for sidewalls made of novel vinyl-cis-polybutadiene, which has a small die swell and excellent extrusion processability and which is capable of improving low fuel consumption with respect to a vulcanizate thereof.

Also, the invention relates to a tire for passenger automobile using, as a cap tread rubber, a rubber composition having a small die swell and excellent extrusion processability while keeping a high elastic modulus.

Also, an object of the invention relates to a rubber composition for large-sized vehicle using, as a cap tread rubber, a rubber composition having a small die swell and excellent extrusion processability while keeping a high elastic modulus.

Also, the invention relates to a high-hardness compounded rubber composition and more specifically, to a rubber composition suitable for beads and chafers of tire, which has improved dimensional stability (die swell is small) at the time of extrusion processing or durability while keeping a high hardness.

Also the invention relates to a rubber composition for base treads and more specifically, to a rubber composition for base treads made of novel vinyl-cispolybutadiene capable of making both internal exothermic characteristics of a tire and extrusion processability compatible with each other in a good balance.

Also, the invention relates to a rubber composition for internal members of tire such as tire coating rubbers inclusive of carcasses and belts, which has a small die swell, excellent extrusion processability and satisfactory adhesiveness to metals.

The rubber composition of the invention can be used for tire members such as tire sidewalls, cap treads, side reinforcing layers of run flat tire, carcasses, belts, chafers, base treads, beads, stiffners, and inner liners; industrial products such as rubber vibration insulators, hoses, belts, rubber rolls, rubber coolers, and shoe sole rubbers; and other composites, adhesives, plastic modifiers, and so on.

Background Art

[0002]

Polybutadiene has a so-called microstructure in which a binding segment formed by polymerization at the 1,4-positions (1,4-structure) and a binding segment formed by polymerization at the 1,2-positions (1,2-structure) are copresent in the molecular chain. The 1,4-structure is further classified into a cis-structure and a transstructure. On the other hand, the 1,2-structure takes a structure containing a vinyl group as a side chain. [0003]

Hitherto, a production process of a vinyl-cispolybutadiene rubber composition has been carried out in an aromatic hydrocarbon-based solvent such as benzene, toluene, and xylene. When such a solvent is used, since the viscosity of a polymerization solution is high, there were problems in stirring, heat transfer, delivery, and so on, and excessive energy was required for recovering the solvent.

[0004]

As the foregoing production process, there is known a process in which 1,3-butadiene is subjected to cis-1,4polymerization in the foregoing inert organic solvent by using a catalyst obtainable from water, a soluble cobalt compound and an organoaluminum chloridehalide represented by the general formula, AlR_nX_{3-n} (wherein R represents an alkyl group having from 1 to 6 carbon atoms, a phenyl group, or a cycloalkyl group; X represents a halogen atom; and n represents a numeral of from 1.5 to 2) to produce BR, and 1,3-butadiene is then subjected to syndiotactic 1,2polymerization (hereinafter abbreviated as polymerization") in the presence of a catalyst obtainable from a soluble cobalt compound, an organoaluminum compound represented by the general formula, AlR3 (wherein R represents an alkyl group having from 1 to 6 carbon atoms,

a phenyl group or a cycloalkyl group) and carbon disulfide by adding or not adding 1,3-butadiene and/or the foregoing solvent to this polymerization system (see, for example, JP-B-49-17666 (Patent Document 1) and JP-B-49-17667 (Patent Document 2)).

[0005]

Furthermore, for example, JP-B-62-171 (Patent Document 3), JP-B-63-36324 (Patent Document 4), JP-B-2-37927 (Patent Document 5), JP-B-2-38081 (Patent Document 6) and JP-B-36366 (Patent Document 7) describe a process in which the production is carried out by subjecting 1,3-butadiene to cis-1,4-polymerization in the presence or absence of carbon disulfide, or after the production, 1,3-butadiene and carbon disulfide are separated and recovered, thereby circulating 1,3-butadiene not substantially containing carbon disulfide or the foregoing inert organic solvent. In addition, JP-B-4-48815 (Patent Document 8) describes a rubber composition having excellent tensile stress and flex crack growth resistance, in which a compounded material thereof has a small die swell ratio and a vulcanizate thereof is suitable as a sidewall of tire.

[0006]

Furthermore, JP-A-2000-44633 (Patent Document describes a process in which the production is carried out in an inert organic solvent containing, as the major component, a C4 fraction such as n-butane, cis-2-butene, trans-2-butene, and butene-1. It is described that in this process, 1,2-polybutadiene contained in the composition is a short fiber crystal, and the distribution of the major axis length of the short fiber crystal is such that 98 % or more of the fiber length is less than 0.6 µm and 70 % or more thereof is less than 0.2 µm. Also, it is described that in the resulting rubber composition, the moldability, tensile stress, tensile strength and flex crack growth resistance of a cis-1,4-polybutadiene rubber (hereinafter abbreviated as "BR") are improved. However, rubber compositions in which various characteristics are improved were demanded depending upon the utility. [0007]

Patent Document 1: JP-B-49-17666
Patent Document 2: JP-B-49-17667
Patent Document 3: JP-B-62-171
Patent Document 4: JP-B-63-36324
Patent Document 5: JP-B-2-37927
Patent Document 6: JP-B-2-38081
Patent Document 7: JP-B-363566
Patent Document 8: JP-B-4-48815
Patent Document 8: JP-B-4-48815
Patent Document 9: JP-A-2000-44633

[8000]

In general, with respect to the performance of automobile tire, there are required excellent wet skid properties as a braking characteristic and excellent rolling resistance (tan δ) or abrasion resistance as a fuel consumption characteristic. However, it is known that these characteristics are in an antinomy relation with each other. In recent years, though there has been made a proposal to compound silica having excellent wet skid properties, thereby highly balancing the foregoing characteristics, such is not satisfactory yet. It is known that though the silica compounding is excellent in the wet skid properties and fuel consumption properties, abrasion resistance and processability are lowered. While the abrasion resistance is improved by the use of high cis-BR, there is some possibility that the wet skid properties are lowered, and therefore, improvements were demanded.

[0009]
Furthermore, in general, it is known that a proportion of the tire rolling resistance occupying in the running resistance which largely affects the fuel consumption of an automobile is large and that the affect due to an energy loss of sidewall rubber is relatively large. For that reason, rubbers having a low dissipation factor such as natural rubbers, isoprene rubbers, butadiene rubbers, and mixtures thereof have hitherto been used in the sidewall part. However, it is required that the rolling resistance of tire is further improved.

[0010]

As a method for reducing the dissipation factor of the sidewall rubber, the use of carbon black having low reinforcing properties or a reduction of the compounding amount of carbon black may be thought. However, there is encountered such a problem that the die swell at the time of extrusion processing becomes large, and therefore, it is difficult to make the thickness of the sidewall member thin or to realize an enhancement of uniformity of tire. For that reason, there is demanded a method for realizing low fuel consumption while keeping a small die swell with respect to an extrudate thereof.

[0011]

Furthermore, in general, tires are required to have excellent drivability and durability and so on. In particular, from the standpoint of safety, the tires are required to have excellent wet skid resistance on a wet road surface. Moreover, on the basis of social requirements for realizing conservation of natural resources in recent years, tires having a low rolling

resistance, namely tires with a low energy loss, are being researched and developed. While the energy loss of tire as consumed by free rotation varies depending upon the tire structure, etc., about a half of the whole energy is consumed in the tread part. Accordingly, if the energy loss of the tread rubber is reduced, a tire with a low energy loss at the time of rolling is obtained.

[0012]

Then, it is attempted to modify the tread rubber such that the energy loss is low. However, such modification of the rubber tends to lower the wet skid properties. Since an improvement of the rolling resistance and an improvement of the wet skid properties generally contradict to each other, various designs for the improvements in the tire structure are attempted in order to make them compatible with each other. As one of the designs, there is enumerated a method for forming a tread into two layers consisting of a cap tread and a base tread. That is, this method is to form the tread into two layers of a cap tread with excellent wet skid properties and a base tread with a low energy loss, thereby not only enhancing the wet skid properties of tire as a whole but also lowering the energy loss.

The rubber for cap tread is required to have a high elastic modulus and molding stability derived from the abrasion resistance and high-speed running properties in addition to the wet skid properties. As a method for obtaining a rubber with a high elastic modulus, various methods have hitherto been attempted. The method for compounding a large amount of carbon black is not preferable because unification of rubbers is poor in the processing step, an electrical power load increases at the time of kneading or extrusion, and ML of the compounded material becomes large, whereby difficulty is accompanied at the time of extrusion molding. The method for compounding a large amount of sulfur involves such defects that sulfur causes blooming and that the crack growth becomes fast due to an increase of the degree of crosslinking. With respect to the addition of thermosetting resin, since the thermosetting resin is low in compatibility with usually used natural rubbers or diene-based rubbers, when a large amount of the thermosetting resin is compounded, satisfactory dispersion is hardly obtained. Furthermore, since this mixed compound is hard even in an unvulcanized state, the load may become large in kneading or extrusion, or the molding processability of tire may be deteriorated. In a method for merely blending and compounding single fibers, since binding between short fibers and the rubber is insufficient, creep may become large, or fatigue life may be lowered.

[0013]

As the rubber for base tread, a rubber with low exothermic characteristics is required. Examples of rubbers with low exothermic characteristics include natural rubbers, isoprene rubbers, and cis-1,4-polybutadiene rubbers, and compounded materials of such a rubber with carbon black are used. In order to reveal low exothermic characteristics by other materials than rubbers, it may be thought to use carbon black having a large particle size and low reinforcing properties or to lower the compounding amount of carbon black. However, these methods involve such problems that the elastic modulus or fatique resistance of rubber is lowered and that the die swell becomes large. Furthermore, there is a method for revealing low exothermic characteristics by compounding short fibers of nylon or vinylon, thereby making the rubber have a high elastic modulus. However, since these short fibers are insufficient in adhesion to rubbers, there is encountered a problem that the fatigue life is short.

On the other hand, when a cap/base system is employed in passenger automobile tires, at the time of co-extrusion of a cap tread and a base tread, there is caused a problem that an extrudate is warped because of a difference in die swell between the both members. If the die swell of the base tread rubber is made small, this problem is liable to be solved. If a large amount of carbon black with high reinforcing properties is compounded, though the die swell become small, the heat generation becomes large. Thus, a method which is satisfied with the die swell and low heat generation at the same time is desired.

[0014]

Furthermore, as rubber members to be disposed in the surroundings of tire bead, high-hardness rubbers are used. Examples thereof include a rubber composition having an increased compounding amount of carbon black and a rubber composition having a novolak-type phenol resin compounded therein (see JP-B-57-30856) and a rubber composition having short fibers and an olefin-based resin compounded therein (see JP-A7-315014).

[0015]

With respect to a method for obtaining a high-hardness rubber composition, it is general to increase carbon black or to compound a fiber, a resin, etc. However, in all of these methods, there may be the case where the heat generation at the time of repeated deformation becomes

large, whereby the durability and fatigue resistance are lowered. Thus, improvements are required. [0016]

Furthermore, in general, in radial tires, a steel cord is used in view of high-speed durability and high-speed drivability. In the case of using a steel cord, a very large strain is liable to be gathered in a rubber in the vicinity of the steel cord at the time of running of tire. Accordingly, the rubber for steel cord is required to have a high elastic modulus and excellent adhesiveness to metals. Even in radial tires or bias tires using an organic fiber cord, rubbers having a high elastic modulus are preferable as the rubber for cord from the viewpoint of durability.

[0017]

As a method for obtaining a rubber with a high elastic modulus, various methods have hitherto been attempted. The method for compounding a large amount of carbon black is not preferable because unification of rubbers is poor in the processing step, an electrical power load increases at the time of kneading or extrusion, and ML of the compounded material becomes large, whereby difficulty is accompanied at the time of extrusion molding. The method for compounding a large amount of sulfur involves such defects that sulfur causes blooming and that the crack growth becomes fast due to an increase of the degree of crosslinking. With respect to the addition of a thermosetting resin, since the thermosetting resin is low in compatibility with natural rubbers or diene-based rubbers as usually used as a cord coating rubber, dispersion failure likely occurs, and crack resistance is deteriorated. Furthermore, conventionally known rubber compositions for tire cord coating are small in green strength. Thus, materials having a large green strength are required in view of molding processability.

Disclosure of the Invention

Problems that the Invention is to Solve

[0018]

An object of the invention is to obtain a silica compounded rubber composition for tire having excellent extrusion processability and having satisfactory wet skid properties and abrasion resistance while keeping a high elastic modulus.

Also, an object of the invention is to obtain a rubber composition for sidewall having low fuel consumption with respect to a vulcanizate thereof and having a small die swell.

Also, an object of the invention is to obtain a tire

for passenger automobile having excellent extrusion molding processability and having good high-speed running properties, grip properties on a wet road surface and abrasion resistance by using a rubber composition for cap tread having excellent extrusion processability while keeping a high elastic modulus.

Also, an object of the invention is to obtain a rubber composition for large-sized vehicle having excellent moldability and having good high-speed running properties, wet skid properties and abrasion resistance by using a rubber composition for cap tread having a small die swell and excellent extrusion processability while keeping a high elastic modulus.

Also, an object of the invention is to provide a rubber composition capable of improving dimensional stability at the time of extrusion processing and durability with respect to a tire thereof while keeping a high hardness.

Also, an object of the invention is to obtain a rubber composition for base tread having low exothermic characteristics with respect to a vulcanizate thereof and having a small die swell.

Also, an object of the invention is to obtain a rubber composition for tire cord coating of carcasses, belts, beads, etc. having a small die swell, a large green strength and excellent molding processability and having a large elastic modulus with respect to a vulcanizate thereof.

Means for Solving the Problems [0019]

(1) The invention is concerned with a rubber composition which is a silica compounded rubber composition for tire comprising 100 parts by weight of a rubber component made of (a) from 20 to 80 % by weight of a vinyl-cispolybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a high-molecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one member selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, polybutadiene and derivatives thereof and (b) from 80 to 20 % by weight of a diene-based rubber other than (a); and (c) from 40 to 100 parts by weight of a rubber reinforcing agent containing 40 % or more of silica, which is characterized in that the 1,2-polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cispolybutadiene rubber which is a matrix component of the

subject vinyl-cis-polybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the high-molecular substance.
[0020]

- (2) Also, the invention is concerned with a rubber composition which is a rubber composition for sidewall comprising 100 parts by weight of a rubber component made of (a) from 20 to 80 % by weight of a vinyl-cispolybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a high-molecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one member selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, polybutadiene and derivatives thereof and (b) from 80 to 20 % by weight of a diene-based rubber other than (a); and (c) from 25 to 60 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2-polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which is a matrix component of the subject vinyl-cis-polybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the high-molecular substance. [0021]
- (3) Also, the invention is concerned with a rubber composition which is a rubber composition for passenger automobile tire comprising 100 parts by weight of a rubber component made of (a) from 10 to 50 % by weight of a vinylcis-polybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a highmolecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, liquid polybutadiene and derivatives thereof, (e) from 30 to 70 % by weight of a styrene-butadiene rubber, and (b) from 0 to 60 % by weight of a diene-based rubber other than (a) and (e); and (d) from 40 to 100 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2-polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which is a matrix component of the subject vinyl-cispolybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the high-molecular substance.

[0022]

- (4) Also, the invention is concerned with a rubber composition which is a rubber composition for large-sized vehicle tire comprising 100 parts by weight of a rubber component made of (a) from 10 to 60 % by weight of a vinylcis-polybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a highmolecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, liquid polybutadiene and derivatives thereof and (b) from 90 to 40 % by weight of a diene-based rubber other than (a); and (c) from 45 to 70 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which is a matrix component of the subject vinyl-cis-polybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the highmolecular substance. [0023]
- (5) Also, the invention is concerned with a rubber composition which is a high-hardness compounded rubber composition comprising 100 parts by weight of a rubber component made of (a) from 20 to 80 % by weight of a vinylcis-polybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a highmolecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one selected from polyisoprene, crvstalline polybutadiene having a melting point of not higher than 150 °C, liquid polybutadiene and derivatives thereof and (b) from 80 to 20 % by weight of a diene-based rubber other than (a); and (c) from 60 to 100 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2-polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which matrix component of the subject vinyl-cispolybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the high-molecular substance.
- (6) Also, the invention is concerned with a rubber composition which is a rubber composition for base tread

[0024]

comprising 100 parts by weight of a rubber component made of (a) from 20 to 80 % by weight of a vinyl-cispolybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a high-molecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one member selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, polybutadiene and derivatives thereof and (b) from 80 to 20 % by weight of a diene-based rubber other than (a); and (c) from 22 to 55 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2-polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which is a matrix component of the subject vinyl-cis-polybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the high-molecular substance. [0025]

Also, the invention is concerned with a rubber (7) composition which is a rubber composition for tire cord coating comprising 100 parts by weight of a rubber component made of (a) from 10 to 60 % by weight of a vinylcis-polybutadiene rubber containing 1,2-polybutadiene having a melting point of 170 °C or higher and a highmolecular substance having at least one unsaturated double bond per a repeating unit and comprising at least one crystalline member selected from polyisoprene, polybutadiene having a melting point of not higher than 150 °C, liquid polybutadiene and derivatives thereof and (b) from 90 to 40 % by weight of a diene-based rubber other than (a); and (c) from 30 to 80 parts by weight of a rubber reinforcing agent, which is characterized in that the 1,2polybutadiene is dispersed in a short crystalline fiber state and the high-molecular substance is dispersed in a granular state in the cis-polybutadiene rubber which is a matrix component of the subject vinyl-cis-polybutadiene rubber (a); and that the short crystalline fiber of the 1,2-polybutadiene is dispersed in particles of the highmolecular substance.

[0026]

(8) The invention is concerned with the rubber composition as set forth above in any one of (1) to (7), which is characterized in that the subject vinyl-cis-polybutadiene rubber (a) is produced by a process for producing a vinyl-cis-polybutadiene rubber by subjecting 1,3-butadiene to cis-1,4-polymerization in a hydrocarbon-based solvent by

using a cis-1,4-polymerization catalyst, subsequently making a 1,2-polymerization catalyst copresent in the resulting polymerization reaction mixture to subject the 1,3-butadiene to 1,2-polymerization, thereby forming 1,2polybutadiene having a melting point of 170 °C or higher. and then separating, recovering and obtaining a vinyl-cispolybutadiene rubber as formed from the resulting polymerization reaction mixture, which is characterizing by including a step for adding a high-molecular substance having at least one unsaturated double bond per a repeating unit within the production system of the vinyl-cispolybutadiene rubber.

[0027]

(9) Also, the invention is concerned with the rubber composition as set forth above in any one of (1) to (8), which is characterized in that in the production step of the vinyl-cis-polybutadiene rubber (a), the unsaturated high-molecular substance is contained in the range of from 0.01 to 50 % by mass based on the total sum of the crystalline fiber of 1,2-polybutadiene and the cis-polybutadiene rubber.

[0028]

(10) Also, the invention is concerned with the rubber composition as set forth above in any one of (1) to (9), which is characterized in that the short fiber of 1,2-polybutadiene in the subject vinyl-cis-polybutadiene rubber (a) is also dispersed in the cis-polybutadiene rubber as the matrix component without being contained in the particles of the high-molecular substance; that the major axis length of the short crystalline fiber dispersed in the subject matrix is in the range of from 0.2 to 1,000 µm; and that the major axis length of the short crystalline fiber of the 1,2-polybutadiene dispersed in the particles of the subject high-molecular substance is in the range of from 0.01 to 0.5 µm.

[0029]

- (11) Also, the invention is concerned with the rubber composition as set forth above in any one of (1) to (10), which is characterized that the subject vinyl-cispolybutadiene rubber (a) has the following characteristics:
- (1) the cis-polybutadiene rubber which is the matrix component of the subject vinyl-cis-polybutadiene rubber has a Mooney viscosity in the range of from 10 to 50;
- (2) the cis-polybutadiene rubber which is the matrix component of the subject vinyl-cis-polybutadiene rubber has a viscosity in a toluene solution at 25 °C in the range of from 10 to 150;

- (3) the cis-polybutadiene rubber which is the matrix component of the subject vinyl-cis-polybutadiene rubber has an [n] in the range of from 1.0 to 5.0;
- (4) the cis-polybutadiene rubber which is the matrix component of the subject vinyl-cis-polybutadiene rubber has a 1,4-cis-structure content in the range of 80 % ore more;
- (5) the 1,2-polybutadiene and the high-molecular substance are dispersed in a physically and/or chemically adsorbed state in the cis-polybutadiene rubber which is the matrix component of the subject vinyl-cis-polybutadiene rubber; and
- (6) the high-molecular substance in the subject vinylcis-polybutadiene rubber is a boiling n-hexane insoluble matter.

[0030]

(12) Also, the invention is concerned with the rubber composition as set forth above in (1), which is characterized that the diene-based rubber (b) other than (a) is a natural rubber and/or polyisoprene and/or a stvrene-butadiene rubber.

[0031]

(13) Also, the invention is concerned with the rubber composition as set forth above in any one of (2) to (7), which is characterized that the diene-based rubber (b) other than (a) is a natural rubber and/or polyisoprene.

[0032]

(14) Also, the invention is concerned with the rubber composition as set forth above in any one of (2) to (7), which is characterized that the rubber reinforcing agent is carbon black.

Advantages of the Invention

[0033]

The silica compounded rubber composition for tire which is used in the invention has a small die swell and excellent extrusion processability and moldability while keeping high wet skid properties, improves the workability of tire manufacture and has excellent abrasion resistance and low fuel consumption.

The rubber composition for sidewall of the invention has low fuel consumption with respective to a vulcanizate thereof and has a small die swell.

The rubber composition for passenger automobile tire according to the invention is able to realize a high elastic modulus and high abrasion resistance while keeping wet skid properties.

The rubber composition for large-sized vehicle tire according to the invention has a small die swell and excellent extrusion processability while keeping a high

elastic modulus and is able to be highly balanced between wet skid properties and abrasion resistance.

The high-hardness compounded rubber composition according to the invention improves dimensional stability at the time of processing and durability of tire at the same time while keeping a high hardness and is able to make the both performances compatible with each in a good balance.

The rubber composition for base tread according to the invention improves die swell and low fuel consumption at the same time and is able to make the both performances compatible with each in a good balance.

The rubber composition for tire cord coating according to the invention has a small die swell, large green strength and excellent extrusion processability and moldability while keeping a high elastic modulus, has excellent adhesiveness to metals and is able to be highly balanced among the respective characteristics. Brief Description of the Drawings

[Fig. 1] shows an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene having a melting point of 170 $^{\circ}$ C or higher and a fine particle 3 of an unsaturated high-molecular substance are individually dispersed in a matrix 1

[Fig. 2] shows an embodiment in which a fine particle 3 of an unsaturated high-molecular substance is dispersed in a matrix 1 in a state that it is attached to a crystalline fiber 2 of 1,2-polybutadiene.

[Fig. 3] shows an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene is dispersed in a matrix 1 in a state that it is attached to a fine particle 3 of an unsaturated high-molecular substance.

[Fig. 4] shows an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene is dispersed in a matrix 1 in a state that it is included and dispersed in a fine particle 3 of an unsaturated high-molecular substance. Best Modes for Carrying Out the Invention

[0035]

In general, the vinyl-cis-polybutadiene rubber (a) of the invention is constructed as follows. That is, it is composed of (1) from 1 to 50 parts by mass of 1,2-polybutadiene having a melting point of 170 °C or higher, (2) 100 parts by mass of a cis-polybutadiene rubber, and (3) from 0.01 to 50 % by mass, based on the whole amount of the foregoing (1) and (2), of an unsaturated high-molecular substance. Furthermore, the 1,2-polybutadiene having a melting point of 170 °C or higher which is the component

(1) forms a crystalline fiber in a short fiber state in which a minor axis length of average monodispersed fiber crystals is not more than 0.2 µm, an aspect ratio is not more than 10, and the number of average monodispersed fiber crystals is 10 or more.

[0036]

It is desired that the crystalline fiber of 1,2polybutadiene which is the foregoing component (1) is in a short fiber state in which a minor axis length of average monodispersed fiber crystals is not more than 0.2 µm, and preferably not more than 0.1 µm, an aspect ratio is not more than 10, and preferably not more than 8, and the number of average monodispersed fiber crystals is 10 or more, and preferably 15 or more, and has a melting point of 170 °C or higher, and preferably from 190 to 220 °C.

[0037]

Then, in the vinyl-cis-polybutadiene rubber (a) of the invention, the 1,2-polybutadiene having a melting point of 170 °C or higher which is the foregoing component (1) is present in a short crystalline fiber state and the unsaturated high-molecular substance which is the foregoing component (3) is present in a granular state in the cispolybutadiene which is the foregoing component (2) as a matrix component. Furthermore, the particle of the unsaturated high-molecular substance as dispersed in the cis-polybutadiene which is the foregoing component (2) as a matrix component has a major axis size in the range of from 0.2 to 1,000 μm , and the short crystalline fiber of the 1,2-polybutadiene as dispersed in the particles of the subject high-molecular substance has a major axis length in the range of from 0.01 to 0.5 μm .

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It is desired that the cis-polybutadiene rubber which foregoing component (2) has the following characteristic. That is, the Mooney viscosity (ML1+4 at 100 °C, hereinafter abbreviated as "ML") is preferably from 10 to 50, and more preferably from 10 to 40. In this way, there are brought such effects that the workability at the time of compounding is improved and the dispersibility of the foregoing component (1) into the component (2) is Furthermore, it is desired that the cispolybutadiene rubber which is the component (2) has the following characteristics. That is, it is desired that the viscosity in a toluene solution (centipoises at 25 °C, hereinafter abbreviated as "T-cp") is preferably from 10 to 150, and more preferably from 10 to 100; and that $[\eta]$

(inherent viscosity) is from 1.0 to 5.0, and preferably from 1.0 to 4.0. Furthermore, it is desired that 1,4-cisstructure content is 80 % or more, and preferably 90 % or more; and that a gel matter is not substantially contained. Here, what a gel matter is not substantially contained means that a toluene insoluble matter is not more than 0.5 % by mass.

100391

The "toluene insoluble matter" as referred to herein means a gel matter as attached to a 200-mesh wire net after charging 10 g of a sample rubber and 400 mL of toluene in an Erlenmeyer flask, completely dissolving at RT (25 °C) and then filtering the solution by using a filter on which the wire net is placed. The foregoing proportion refers to a value as defined in terms of percentage to the sample rubber, which is obtained by drying in vacuo the wire net having a gel attached thereonto and measuring the attachment amount.

[0040]

Furthermore, the $[\eta]$ (inherent viscosity) is a value $[\eta]$ as determined according to the following expressions after charging 0.1 g of a sample rubber and 100 mL of toluene in an Erlenmeyer flask, completely dissolving at 30 °C, charging 10 mL of the solution in a Cannon-Fenske dynamic viscometer in a constant-temperature water tank controlled at 30 °C and measuring a drop time (T) of the solution.

 $\eta sp = T/T_0 - 1$ (T₀: drop time of only toluene) $\eta sp/c = [\eta] + k'[\eta]^2c$

(η sp: specific viscosity, k': Huggins' constant (0.37), c: sample concentration (g/mL) [0041]

It is desired that with respect to the proportion of the 1,2-polybutadiene crystalline fiber which is the foregoing component (1) to the cis-polybutadiene rubber which is the component (2), the proportion of the 1,2polybutadiene crystalline fiber which is the foregoing component (1) is from 1 to 50 parts by mass, and preferably from 1 to 30 parts by mass based on 100 parts by mass of the cis-polybutadiene rubber which is the component (2) as described previously. When the proportion falls within the foregoing range, it is possible to avoid inconveniences that in the case where the proportion is large as more than 50 parts by mass, the short fiber crystal of the 1,2-polybutadiene crystalline fiber in the cis-polybutadiene rubber is liable to become large to have poor dispersibility and that in the case where the

proportion is small as less than 1 part by mass, the reinforcing properties due to the short fiber crystal are Accordingly, the problems that the elastic modulus, extrusion processability, moldability and the like as the characteristic features are hardly revealed and that the processability is deteriorated are hardly caused, and therefore, such is preferable. Furthermore, it is desired that the proportion of the unsaturated high-molecular substance which is the component (3) is from 0.01 to 50 % by mass, and preferably from 0.01 to 30 % by mass based on the vinyl-cis-polybutadiene rubber as described previously. What the proportion falls within the foregoing range is preferable in view of an improvement of the dispersibility due to coagulation of the 1,2-polybutadiene crystalline fiber which is the foregoing component (1) and inhibition in lowering of accompanied various physical properties which are derived from the vinyl-cis-polybutadiene rubber, and so on.

[0042]

For example, the foregoing vinyl-cis-polybutadiene rubber is suitably obtained by the following production process.

[0043]

In general, in the production of the vinyl-cispolybutadiene rubber of the invention, the polymerization of 1,3-butadiene is carried out by using a hydrocarbon-based solvent. As this hydrocarbon-based solvent, a hydrocarbon-based solvent having a solubility parameter (hereinafter abbreviated as "SP value") of not more than 9.0 is preferable, and a hydrocarbon-based solvent having a solubility parameter of not more than 8.4 is more preferable. Examples of the hydrocarbon-based solvent having a solubility parameter of not more than 9.0 include aliphatic hydrocarbons and alicyclic hydrocarbons such as n-hexane (SP value: 7.2), n-pentane (SP value: 7.5), cyclohexane (SP value: 8.1), and n-butane (SP value: 6.6). Of these, cyclohexane or the like is preferable.

[00441

The SP values of these solvents are known by documents such as Rubber Industry Handbook (Fourth Edition, The Society of Rubber Industry, Japan, issued January 20, 1994, page 721).

[0045]

By using a solvent having an SP value of lessnot more
than 9.0, the dispersion state of the short fiber crystal
of the 1,2-polybutadiene crystalline fiber into the cispolybutadiene rubber is formed as expected in the invention

and an excellent die swell characteristic and high elastic modulus and tensile strength are revealed, and therefore, such is preferable.

[0046]

First of all, 1,3-butadiene and the foregoing solvent are mixed, and the water content in the resulting solution is then adjusted. The water content is preferably in the range of from 0.1 to 1.0 mole, and especially preferably from 0.2 to 1.0 mole per mole of the organoaluminum chloride used as the after-mentioned cis-1,4-polymerization catalyst in the foregoing solution. What the water content falls within this range is preferable because the sufficient catalytic activity can be obtained; preferable content of the cis-1,4-structure and the molecular weight can be obtained; the formation of a gel at the time of polymerization can be inhibited so that attachment of the gel to a polymerization tank or the like can be prevented; and in addition, the continuous polymerization time can be prolonged. As a method for adjusting the water content, known methods can be applied. A method for undergoing addition and dispersion by passing through a porous filtering material (see JP-A-4-85304) is also effective.

[0047]

An organoaluminum chloride is added as one of the cish, 4-polymerization catalysts to the solution obtained by adjusting the water content. As the organoaluminum chloride, a compound represented by the general formula, AlR_{n} -n is preferably used. As specific examples thereof, diethylaluminum monochloride, diethylaluminum monochloride, diethylaluminum monochloride, dicyclohexylaluminum monochloride, diphenylaluminum monochloride, diethylaluminum seaquichloride, and the like can be suitably enumerated. The amount of use of the organoaluminum chloride is preferably 0.1 mmoles or more, and more preferably from 0.5 to 50 mmoles per mole of the whole amount of the 1,3-butadiene.

[0048]

Next, a soluble cobalt compound is added as one of other cis-1,4-polymerization catalysts to the mixed solution having an organoaluminum chloride added thereto, thereby subjecting the 1,3-dibutadiene to cis-1,4-polymerization. The soluble cobalt compound is a cobalt compound which is soluble in a hydrocarbon-based solvent to be used or is soluble in liquid 1,3-butadiene, or is able to be uniformly dispersed. Examples thereof include β -diketone complexes of cobalt such as cobalt(II) acetylacetonate and cobalt(III) acetylacetonate; β -keto-

acid ester complexes of cobalt such as a cobalt ethylacetoacetate complex; cobalt salts of an organic carboxylic acid having 6 or more carbon atoms such as cobalt octoate, cobalt naphthenate, and cobalt benzoate; and halogenated cobalt complexes such as a cobalt chloride pyridine complex and a cobalt chloride ethyl alcohol complex. The amount of the soluble cobalt compound to be used is 0.001 mmoles or more, and especially preferably 0.005 mmoles or more based on one mole of 1,3-butadiene. Furthermore, a molar ratio of the organoaluminum chloride to the soluble cobalt compound (Al/Co) is 10 or more, and especially preferably 50 or more. Moreover, in addition to the soluble cobalt compound, an organic carboxylic acid salt of nickel, an organic complex salt of nickel, an organolithium compound, an organic carboxylic acid salt of neodymium, and an organic complex salt of neodymium can be used.

[0049]

The temperature at which the cis-1,4-polymerization is carried out is generally from a temperature exceeding 0 °C to 100 °C, preferably from 10 to 100 °C, preferably from 20 to 100 °C. The polymerization time (average residence time) is preferably in the range of from 10 minutes to 2 hours. It is preferred to carry out the cis-1,4-polymerization such that the polymer concentration after the cis-1,4-polymerization is from 5 to 26 % by mass. As the polymerization tank, a single tank is used or two or more tanks are connected and used. The polymerization is carried out by stirring and mixing the solution within the polymerization tank (polymerizer). As the polymerization tank used for the polymerization, a polymerization tank equipped with a high-viscosity solution stirring unit, for example, a device as described in JP-B-40-2645 can be emploved.

[0050]

In the production of the vinyl-cis-polybutadiene rubber of the invention, at the time of the cis-1,4-polymerization, a known molecular weight modifier, for example, non-conjugated dienes such as cyclooctadiene, allene, and methylallene (1,2-butadiene); and α -olefins such as ethylene, propylene, and butene-1 can be used. Furthermore, in order to further inhibit the formation of a gel at the time of the polymerization, a known gelation preventing agent can be used. Furthermore, in general, the content of the cis-1,4-structure in the polymerization product is 80 % or more, and preferably from 10 to 50, and preferably from 10 to 40. A

gel matter is not substantially contained.

Then, by adding an organoaluminum compound represented by the general formula, AlR3 and carbon disulfide and optionally, the foregoing soluble cobalt compound as 1,2polymerization catalysts to the thus obtained cis-1,4polymerization reaction mixture, the 1,3-butadine is subjected to 1,2-polymerization, thereby producing a vinylcis-polybutadiene rubber. On this occasion, the unreacted 1,3-butadiene may be reacted by adding or not adding 1,3butadiene to the subject polymerization reaction mixture. Suitable examples of the organoaluminum compound represented by the general formula, AlR₃ include trimethylaluminum, triethylaluminum, triisobutylaluminum, tri-n-hexylaluminum, and triphenylaluminum. The amount of the organoaluminum compound is 0.1 mmoles or more, and especially from 0.5 to 50 mmoles-or-more per mole of the 1,3-butadiene. Though the carbon disulfide is not particularly limited, water-free carbon disulfide is preferable. The concentration of the carbon disulfide is not more than 20 mmoles/L, and especially preferably from 0.01 to 10 mmoles/L. As a replacement of the carbon disulfide, known phenyl isothiocyanate and xanthogenic acid compounds may be used.

[0052]

The temperature at which the 1,2-polymerization is carried out is generally from 0 to 100 °C, preferably from 10 to 100 °C, and more preferably from 20 to 100 °C. the 1,2-polymerization, by adding from 1 to 50 parts by mass, and preferably from 1 to 20 parts by mass, based on 100 parts by mass of the foregoing cis-1,4-polymerization reaction mixture, of 1,3-butadiene to the polymerization system, the yield of 1,2-polybutadiene at the time of the 1,2-polyerization can be increased. The polymerization time (average residence time) is preferably in the range of from 10 minutes to 2 hours. It is preferred to carry out the 1,2-polymerization such that the polymer concentration after the 1,2-polymerization is from 9 to 29 % by mass. As the polymerization tank, a single tank is used or two or more tanks are connected and used. The polymerization is carried out by stirring and mixing the polymerization solution within the polymerization tank (polymerizer). With respect to the polymerization tank which is used for the 1,2-polymerization, since the viscosity becomes high during the 1,2-polymerization so that the polymer is liable to attach thereto, a polymerization tank equipped with a high-viscosity solution stirring unit, for example, a device as described in JP-B-40-2645 can be employed.

[0053]

In the production of the vinyl-cis-polybutadiene rubber of the invention, when the vinvl-cis-polybutadiene rubber is produced by carrying out the cis-1,4polymerization and subsequently the 1.2-polymberization as described previously, there is included a step for adding a high-molecular substance having at least one unsaturated double bond per a repeating unit and comprising at least member selected from polyisoprene, crystalline polybutadiene having a melting point of not higher than 150 °C, liquid polybutadiene and derivatives thereof within the production system of the vinyl-cis-polybutadiene rubber. After the production of a vinyl-cis-polybutadiene rubber, for example, even when such a substance is added at the time of compounding, the effects of the invention of this application are not obtained. With respect to the addition of this unsaturated high-molecular substance within the production system, the addition in the polymerization reaction mixture is preferably carried out at an arbitrary point of time of from the cis-1,4-polymerization to the 1,2-polymerization, and more preferably at the time of the 1,2-polymerization.

[0054]

As the foregoing unsaturated high-molecular substance, at least one member selected from polyisoprene, crystalline polybutadiene having a melting point of lower than 170 $^{\circ}\mathrm{C}$, linear polybutadiene, and derivatives thereof is preferable.

[0055]

Examples of the polyisoprene include usually synthetic polyisoprene (for example, cis-1,4-polyisoprene in which the content of the cis-structure is 90 % or more), liquid polyisoprene, trans-polyisoprene, and other modified polyisoprene.

[0056]

The crystalline polybutadiene having a melting point of lower than 170 °C is preferably crystalline polybutadiene having a melting point of from 0 to 150 °C, and examples thereof include low-melting point 1,2-polybutadiene and trans-polybutadiene.

[0057]

Examples of the liquid polybutadiene include extremely low-molecular polybutadiene having an inherent viscosity $[\eta]$ of not more than 1.

[0058]

Furthermore, examples of these derivatives include an isoprene/isobutylene copolymer, an isoprene/styrene

copolymer, a styrene/isoprene/styrene block copolymer, liquid epoxidized polybutadiene, liquid carboxyl-modified polybutadiene, and hydrogenation products of these derivatives.

[0059]

Of the foregoing respective unsaturated high-molecular substances, isoprene/styrene block copolymer, and 1,2-polybutadiene having a melting point of from 70 to 110 °C are preferably used. Further, the foregoing respective unsaturated high-molecular substances can be used singly in admixture of two or more kinds thereof.

100601

As described previously, when the foregoing unsaturated high-molecular substance is added, in the resulting vinyl-cis-polybutadiene rubber, the dispersibility of 1,2-polybutadiene having a melting point of 170 °C or higher into the cis-polybutadiene rubber as a matrix component is remarkably enhanced due to a compatible effect of the unsaturated high-molecular substance. As a result, the characteristics of the resulting vinyl-cis-polybutadiene rubber become excellent.

[0061]

The amount of addition of the unsaturated high-molecular substance is preferably in the range of from 0.01 to 50 % by mass, and more preferably from 0.01 to 30 % by mass based on the vinyl-cis-polybutadiene rubber to be obtained. Furthermore, with respect to the addition at any point of time, after the addition, stirring is preferably carried out for from 10 minutes to 3 hours, and more preferably from 10 minutes to 30 minutes.

[0062]

After the polymerization reaction has reached a predetermined rate of polymerization, a known antioxidant can be added according to the usual way. Representative of the antioxidant include phenol-based examples antioxidants such as 2,6-di-t-butyl-p-cresol phosphorus-based antioxidants such as trinonylphenyl phosphite (TNP); and sulfur-based antioxidants such as 4,6bis(octylthiomethyl)-o-cresol and dilauryl-3,3'thiodipropionate (TPL). The antioxidant may be used singly or in combination of two or more kinds thereof. The amount of addition of the antioxidant is from 0.001 to 5 parts by mass based on 100 parts by mass of vinyl-cis-polybutadiene rubber. Next, a short stop agent is added to the polymerization system, thereby stopping the polymerization. This is carried out by a method which is known by itself, for example, a method in which after completion of the polymerization reaction, the polymerization reaction mixture is fed into a short stop tank and a large amount of a polar solvent such as alcohols such as methanol and ethanol and water is charged in this polymerization reaction mixture; and a method in which an inorganic acid such as hydrochloric acid and sulfuric acid, an organic acid such as acetic acid and benzoic acid, or a hydrogen chloride gas is introduced into the polymerization reaction mixture. Next, the formed vinyl-cis-polybutadiene rubber is separated, recovered, washed and dried to obtain the desired vinyl-cis-polybutadiene rubber according to the usual way.

[0063]

In general, in the thus obtained vinvl-cispolybutadiene rubber of the invention, the ratio of the respective components, namely the ratio of the 1,2polybutadiene having a melting point of 170 °C or higher, the cis-polybutadiene rubber and the unsaturated highsubstance is as described previously. Furthermore, in the microstructure of the cis-polybutadiene rubber, 80 % or more thereof is occupied by cis-1,4polybutadiene, and the remainder consists of trans-1,4polybutadiene and vinyl-1,2-polybutadiene. This cispolybutadiene rubber and the unsaturated high-molecular substance are a boiling n-hexane soluble matter, and the 1,2-polybutadiene having a melting point of 170 °C or higher is a boiling n-hexane insoluble matter (hereinafter abbreviated as "H.I."). In general, this 1,2-polybutadiene having a melt point of 170 °C or higher has a melting point of from 170 to 220 °C and is a crystalline fiber in a short fiber state as described previously. Furthermore, as described previously, the ML of the cis-polybutadiene rubber is from 10 to 50, and preferably from 20 to 40.

[0064]

Furthermore, as described previously, in the vinylcis-polybutadiene rubber of the invention, the 1,2polybutadiene having a melting point of 170 °C or higher and the unsaturated high-molecular substance are uniformly dispersed in the matrix of the cis-polybutadiene rubber. [0065]

In general, in the vinyl-cis-polybutadiene rubber of the invention, as described previously, the 1,2-polybutadiene having a melting point of 170 °C or higher is dispersed as a crystalline fiber. Furthermore, the unsaturated high-molecular substance can be dispersed in an embodiment of every kind in relation to the crystalline fiber of the 1,2-polybutadiene having a melting point of

170 °C or higher. Examples of this dispersion embodiment include an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene having a melting point of 170 °C or higher and a fine particle 3 of an unsaturated highmolecular substance are individually dispersed in a matrix 1 as conceptually shown in Fig. 1; an embodiment in which a fine particle 3 of an unsaturated high-molecular substance is dispersed in a matrix 1 in a state that it is attached to a crystalline fiber 2 of 1,2-polybutadiene conceptually shown in Fig. 2; an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene is dispersed in a matrix 1 in a state that it is attached to a fine particle unsaturated high-molecular substance conceptually shown in Fig. 3; and an embodiment in which a crystalline fiber 2 of 1,2-polybutadiene is dispersed in a matrix 1 in a state that it is included and dispersed in a fine particle 3 of an unsaturated high-molecular substance as conceptually shown in Fig. 4. Embodiments in which two kinds or more of the dispersion states as shown in Figs. 1 to 4 coexist may be employed. In Figs. 1 to 4, 1 stands for a matrix; 2 stands for a crystalline fiber of 1,2polybutadiene having a melting point of 170 °C or higher; and 3 stands for a fine particle of an unsaturated highmolecular substance.

[0066]

the production process of the vinyl-cispolybutadiene rubber of the invention, the 1,3-butadiene and the hydrocarbon-based solvent are separated from a mother liquor of the polymerization reaction mixture which remains after separating and obtaining the formed vinylcis-polybutadiene rubber and which contains the unreacted 1,3-butadiene, the hydrocarbon-based solvent and the carbon disulfide and so on usually by distillation; and the carbon disulfide is separated and removed by an adsorption and separation treatment of carbon disulfide or a separation treatment of a carbon disulfide adduct, thereby recovering the 1,3-butaidene and the hydrocarbon-based solvent which do not. substantially contain carbon disulfide. Furthermore, 1,3-butadiene and the hydrocarbon-based solvent which do not substantially contain carbon disulfide can also be recovered by recovering the three components from the foregoing mother liquor of the polymerization reaction mixture by distillation and then separating and removing the carbon disulfide by the foregoing adsorption and separation or separation treatment of a carbon disulfide adduct. The thus recovered carbon disulfide and hydrocarbon-based solvent are mixed with 1,3-butaduene as newly replenished and then provided for reuse. [0067]

According to the foregoing production process of the vinyl-cis-polybutadiene rubber, it is possible to continuously produce the vinyl-cis-polybutadiene rubber of the invention industrially advantageously with excellent operability of the catalyst component and in a high catalytic efficiency. In particular, it is possible to continuously produce the vinyl-cis-polybutadiene rubber industrially advantageously in a high conversion without causing attachment to an inner wall in the polymerization tank, a stirring blade and other portions where stirring is slow.

[0068]

Then, in order that the thus produced vinyl-cispolybutadiene rubber reveals excellent desired characteristics, it is preferable that the polybutadiene crystalline fiber as dispersed in the vinvlcis-polybutadiene rubber is partially dispersed in a monodispersed state as a fine crystal in the matrix of the cis-polybutadiene rubber and coexists together with the large 1,2-polybutadiene crystalline fiber having a coagulation structure. That is, it is preferable that the monodispersed 1,2-polybutadiene crystalline fiber in the matrix of the cis-polybutadiene rubber is in a short fiber state in which a minor axis length of average monodispersed fiber crystals is not more than 0.2 µm, an aspect ratio is not more than 10, and the number of average monodispersed fiber crystals is 10 or more and has a melting point of 170 °C or higher. Furthermore, in addition to the 1,2polybutadiene crystalline fiber having a melting point of 170 °C or higher, it is preferable that the unsaturated high-molecular substance is dispersed in the matrix of the cis-polybutadiene rubber. It is preferable that this unsaturated high-molecular substance has high affinity with the 1,2-polybutadiene crystalline fiber and is dispersed in a physically or chemically adsorbed state in the vicinity of the subject crystalline fiber in the matrix of the cispolybutadiene rubber (the dispersion embodiments of Figs. 2 to 4). As described previously, when the 1,2-polybutadiene crystalline fiber having a melting point of 170 °C or higher and the unsaturated high-molecular substance coexist and are dispersed in the matrix of the cis-polybutadiene rubber, the foregoing various physical properties become excellent, and therefore, such is preferable.

[0069]

Next, a silica compounded rubber composition for tire

which is used in the invention is formed by compounding (a) the foregoing vinyl-cis-polybutadiene rubber, (b) a dienebased rubber other than (a), and (c) a rubber reinforcing agent containing 40 % or more of silica.

[0070]

Also, a rubber composition for cap tread of passenger automobile tire which is used in the invention is formed by compounding (a) the foregoing vinyl-cis-polybutadiene rubber, (e) a styrene-butadiene rubber, (b) a diene-based rubber other than (a) and (e), and (d) a rubber reinforcing agent.

[00711

Also, a rubber composition for sidewall, a rubber composition for large-sized vehicle tire, a high-hardness compounded rubber composition, a rubber composition for base tread and a rubber composition for tire cord coating which are used in the invention are each formed by compounding (a) the foregoing vinyl-cis-polybutadiene rubber, (b) a diene-based rubber other than (a), and (c) a rubber reinforcing agent. [0072]

Examples of the foregoing diene-based rubber (b) include a high cis-polybutadiene rubber, a low cispolybutadiene rubber (BR), a natural rubber, a polyisoprene rubber, an emulsion polymerized or solution polymerized styrene-butadiene rubber (SBR), an ethylene-propylene-diene rubber (EPDM), a nitrile rubber (NBR), a butvl rubber (IIR), and a chloroprene rubber (CR).

[0073]

Furthermore, derivatives of these rubbers, example, polybutadiene rubbers modified with a tin compound and the foregoing rubbers which are subjected to epoxy modification, silane modification or maleic modification can also be used. These rubbers may be used singly or in combination with two or more kinds thereof. [0074]

As the rubber reinforcing agent which is the component (c) of the invention, carbon black having a particle size of not more than 90 nm and having an oil absorption of dibutyl phthalate (DBP) of 70 mL/100 g or more is enumerated, and examples thereof include FEF, FF, GPF, SAF, ISAF, SRF, and HAF. Furthermore, examples of the silica include silicic acid anhydride by the dry method and hydrated silicic acid and synthetic silicates by the wet method. In addition, examples of the rubber reinforcing agent include, in addition to various carbon blacks, inorganic reinforcing agents such as white carbon, activated calcium carbonate, and ultra-finely divided magnesium silicate; and organic reinforcing agents such as a syndiotactic 1,2-polybutadiene resin, a polyethylene resin, a polypropylene resin, a hi-styrene resin, a phenol resin, lignin, a modified melamine resin, a coumaroneindene resin, and a petroleum resin.

[00751

In the silica compounded rubber composition for tire of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 20 to 80 % by weight of the vinyl-cis-polybutadiene (a) and from 80 to 20 % by weight of the diene-based rubber (b) other than (a) and from 40 to 100 parts by weight of the rubber reinforcing agent containing 40 % or more of silica (c).

When the amount of the foregoing vinvl-cispolybutadiene is less than the foregoing lower limit, the die swell is large, and a rubber composition having low exothermic characteristics with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit, the elastic modulus of a vulcanizate thereof is lowered, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that the moldability of tire is liable to become worse. Furthermore, when the proportion of the rubber falls outside the foregoing range, the abrasion resistance of a vulcanizate thereof and the like are lowered. [0076]

In the rubber composition for sidewall of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 20 to 80 % by weight of the vinyl-cis-polybutadiene (a) and from 80 to 20 % by weight of the diene-based rubber (b) other than (a) and from 25 to 60 parts by weight of the rubber reinforcing agent (c).

When the amount of the foregoing vinyl-cispolybutadiene is less than the foregoing lower limit, the die swell is large, and a rubber composition having low fuel consumption with respect to a vulcanizate thereof is not obtained, whereas when the amount of the foregoing vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the

foregoing lower limit, the die swell becomes large, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that kneading becomes difficult, and therefore, such is not preferable. [0077]

In the rubber composition for passenger automobile tire of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 10 to 50 % by weight of the vinyl-cis-polybutadiene (a), from 30 to 70 % by weight of the styrene-butadiene rubber (b) and from 0 to 60 % by weight of the diene-based rubber (c) other than (a) and (b) and from 40 to 100 parts by weight of the rubber reinforcing agent (d).

When the amount of the foregoing vinyl-cispolybutadiene is less than the foregoing lower limit, a rubber composition having a high elastic modulus with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Moonev viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit, the elastic modulus of a vulcanizate thereof is lowered, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that moldability of tire is liable to become Furthermore, when the proportion of the rubber falls outside the foregoing range, the abrasion resistance of a vulcanizate thereof and the like are lowered. [0078]

In the rubber composition for large-sized vehicle tire of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 10 to 60 % by weight of the vinyl-cis-polybutadiene (a) and from 90 to 40 % by weight of the diene-based rubber (b) other than (a) and from 45 to 70 parts by weight of the rubber reinforcing agent (c).

When the amount of the foregoing vinyl-cispolybutadiene is less than the foregoing lower limit, a rubber composition having a high elastic modulus with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit,

the elastic modulus of a vulcanizate thereof is lowered, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that the moldability of tire is liable to become worse. Furthermore, when the proportion of the rubber falls outside the foregoing range, the elastic modulus of a vulcanizate thereof and the like are lowered. [0079]

In the high-hardness compounded rubber composition of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 20 to 80 % by weight of the vinyl-cis-polybutadiene (a) and from 80 to 20 % by weight of the diene-based rubber (b) other than (a) and from 40 to 100 parts by weight of the rubber reinforcing agent (c).

When the amount of the foregoing vinyl-cispolybutadiene is less than the foregoing lower limit, the die swell is large, and a rubber composition having low exothermic characteristics with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit, the die swell becomes large, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that kneading becomes difficult, and therefore, such is not preferable. 100801

In the rubber composition for base tread of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 20 to 80 % by weight of the vinyl-cis-polybutadiene (a) and from 80 to 20 % by weight of the diene-based rubber (b) other than (a) and from 25 to 55 parts by weight of the rubber reinforcing agent (c).

When the amount of the foregoing vinyl-cispolybutadiene is less than the foregoing lower limit, the die swell is large, and a rubber composition having low exothermic characteristics with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit, the die swell becomes large, whereas

when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that kneading becomes difficult, and therefore, such is not preferable.
[0081]

In the rubber composition for tire cord coating of the invention, the foregoing respective components are compounded so as to satisfy the condition of 100 parts by weight of a rubber component of from 10 to 60 % by weight of the vinyl-cis-polybutadiene (a) and from 90 to 40 % by weight of the diene-based rubber (b) other than (a) and from 30 to 80 parts by weight of the rubber reinforcing agent (c).

amount of the foregoing vinvl-cis-When the polybutadiene is less than the foregoing lower limit, a rubber composition having a high elastic modulus with respect to a vulcanizate thereof is not obtained, whereas when the amount of the vinyl-cis-polybutadiene exceeds the foregoing upper limit, the Mooney viscosity of the composition is excessively high so that the moldability becomes worse. When the amount of the foregoing rubber reinforcing agent is less than the foregoing lower limit, the elastic modulus of a vulcanizate thereof is lowered, whereas when it exceeds the foregoing upper limit, the Mooney viscosity is excessively high so that moldability of tire is liable to become Furthermore, when the proportion of the rubber falls outside the foregoing range, the elastic modulus of a vulcanizate thereof and the like are lowered. adhesiveness to metals is lowered.

[0082]
The rubber composition of the invention is obtained by kneading the foregoing respective components in the usual way by using a Banbury, an open roll, a kneader, a twinscrew kneading machine, etc. The kneading temperature is required to be lower than the melting point of the 1,2-polybutadiene crystal fiber which is contained in the subject vinyl-cis-polybutadiene. When kneading is carried out at a temperature of higher than this melting point of the 1,2-polybutadiene crystal fiber, fine short fibers in the vinyl-cis-polybutadiene are melted and deformed into a spherical particle or the like, and therefore, such is not preferable.

[0083]

In the rubber composition of the invention, if desired, compounding agents which are usually used in the rubber field, such as a vulcanizer, a vulcanization aid, an antioxidant, a filler, a process oil, zinc white, and stearic acid, may be kneaded.

[0084]

As the vulcanizer, known vulcanizers such as sulfur, organic peroxides, resin vulcanizers, and metal oxides such as magnesium oxide are useful.

[0085]

As the vulcanization aid, known vulcanization aids such as aldehydes, ammonias, amines, guanidines, thioureas, thiazoles, thiurams, dithiocarbamates, and xanthates are useful.

[0086]

Examples of the antioxidant include amine/ketone-based antioxidants, imidazole-based antioxidants, amine-based antioxidants, sulfur-based antioxidants, sulfur-based antioxidants, and phosphorus-based antioxidants.

[0087]

Examples of the filler include inorganic fillers such as calcium carbonate, basic magnetic carbonate, clay, litharge, and diatomaceous earth; and organic fillers such regenerated rubbers and powdered rubbers.

[0088]

As the process oil, all of aromatic process oils, naphthene-based process oils, and paraffin-based process oils are useful.

[00891

The silica compounded rubber composition for tire of the invention can be designed to enhance the die swell performance, the abrasion resistant performance and the low fuel consumption performance while keeping the wet skid properties and is suitable for applications such as a tread or sidewall of a tire with a more highly balance of the performances.

The rubber composition for sidewall of the invention has a small die swell and has a low exothermic characteristic with respect to a vulcanizate thereof. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known rubber compositions for sidewall.

The rubber composition for passenger automobile tire of the invention has excellent extrusion molding processability and has satisfactory characteristics in high-speed running properties, grip properties on a wet road surface and abrasion resistance. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall

reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known rubber compositions for cap tread.

The rubber composition for large-sized vehicle tire of the invention has a small die swell and has excellent extrusion processability while keeping a high elastic modulus. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known rubber compositions for cap tread

The high-hardness compounded rubber composition of the invention has a small die swell and has a low exothermic characteristic with respect to a vulcanizate thereof. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known high-hardness compounded rubber compositions.

The rubber composition for base tread of the invention has a small die swell and has a low exothermic characteristic with respect to a vulcanizate thereof. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known rubber compositions for base tread.

The rubber composition for tire cord coating of the invention has a small die swell and a large green strength and has excellent extrusion molding processability and moldability while keeping a high elastic modulus. Thus, it can be used as a tire member of passenger automobile, bus, truck, airplane, run flat tire, etc. in combination with other tire member (for example, cap tread, sidewall, sidewall reinforcing layer, base tread, carcass, belt, and bead) in place of the conventionally known rubber compositions for tire cord coating.

[0090]

Examples based on the invention and Comparative Examples will be specifically described below. In the Examples and Comparative Examples, physical properties of a raw rubber of the resulting vinyl-cis-polybutadiene rubber, physical properties of a compounded material of the resulting rubber composition, and physical properties of a

vulcanizate thereof were measured in the following manners.

(1) Content of 1,2-polybutadiene crystalline fiber:

An extraction residue resulting from subjecting 2 g of a vinyl-cis-polybutadiene rubber to boiling extraction with 200 mL of n-hexane for 4 hours by a Soxhlet extractor was expressed in terms of parts by weight.

(2) Melting point of 1,2-polybutadiene crystalline fiber:

An extraction residue of boiling n-hexane was determined by a peak temperature of the endothermic curve by a differential scanning calorimeter (DSC).

(3) Crystalline fiber form:

The vinyl-cis-polybutadiene rubber was vulcanized with sulfur monochloride and carbon disulfide; a vulcanizate was cut out into an ultra-thin section; and the double bond of the rubber segment of the vinyl-cis-polybutadiene was dyed with an osmium tetrachloride vapor and determined by observation by a transmission electron microscope.

(4) Microstructure of rubber segment in vinyl-cispolybutadiene rubber:

An infrared absorption spectral analysis was carried out. The microstructure was calculated from an absorption intensity ratio of cis at $740~\rm cm^{-1}$, trans at $967~\rm cm^{-1}$ and vinyl at $910~\rm cm^{-1}$.

(5) Viscosity in toluene solution of rubber segment in vinyl-cis-polybutadiene rubber:

A viscosity in a 5 % by weight toluene solution at 25 $^{\circ}\mathrm{C}$ was measured and expressed in terms of centipoises (cp).

(6) $[\eta]$ of rubber segment in vinyl-cis-polybutadiene rubber:

A boiling n-hexane soluble matter was collected by drying and measured in a toluene solution at a temperature of 30 °C.

(7) Mooney viscosity:

A value measured at 100 °C according to JIS K6300.

(8) Die swell:

As an estimation for extrusion processability of a compounded material, a ratio of the size of a compounded material and the die orifice size (wherein L/D = 1.5 mm/1.5 mm) was measured and determined at the time of extrusion at 100 $^{\circ}\mathrm{C}$ and at a shear rate of 100 sec^1 by using a processability analyzer (MPT, manufactured by Monsanto).

(9) Tensile elastic modulus:

A tensile elastic modulus M300 was measured according to JIS K6301.

(10) Pico abrasion:

A Pico abrasion index was measured according to ASTM D2228.

(11) Wet skid properties:

This was measured by using a potable wet skid tester and 3M's Safety-Walk (Type B).

(12) Low fuel consumption (low exothermic characteristics or exothermic characteristic):

A heat build-up of a rubber composition was measured in a fixed stress mode with a static load of 196 N under conditions of a temperature of 40 $^{\circ}$ C and a frequency of 10 Hz. The smaller the index, the more excellent the fuel consumption is.

(13) Hardness:

This was measured at room temperature by using a type A durometer according to JIS K6253.

(14) Green modulus:

An unvulcanized rubber was punched out by a No. 3 dumbbell to prepare a specimen, which was then measured at room temperature at a drawing rate of 200 mm/min.

(15) Adhesive strength to metal:

This was measured according to ASTM D2229.

(Production of vinyl-cis-polybutadiene sample 1)

In a stirrer-equipped stainless steel-made reaction tank having an internal volume of 30 L and purged with a nitrogen gas, a solution of 1.6 kg of 1,3-butadiene dissolved in 18 kg of absolute cyclohexane was charged, to which were then added 4 mmoles of cobalt octoate, 84 mmoles diethylaluminum chloride and 70 mmoles of 1,5cyclooctadiene, and the mixture was stirred at 25 °C for 30 minutes, thereby undergoing cis-polymerization. resulting polymer had an ML of 33 and a T-cp of 59 and had a microstructure consisting of 0.9 % of the 1,2-structure, 0.9 % of the trans-1,4-structure and 98.2 % of the cis-1,4-After the cis-polymerization, 5 % by mass structure. (percentage based on the resulting vinyl-cis-polybutadiene rubber) of an unsaturated high-molecular substance consisting of polyisoprene (IR) (ML: 87, cis-1,4-structure: 98 %) was added to the resulting polymerization product solution, and the mixture was stirred at 25 °C for one Immediately thereafter, 90 hour. mmoles triethylaluminum and 50 mmoles of carbon disulfide were added to the polymerization solution, and the mixture was stirred at 25 °C for 60 minutes, thereby undergoing 1,2polymerization. After completion of the polymerization, the polymerization product solution was added to 18 L of methanol containing 1 % by mass of 4.6bis(octylthiomethyl)-o-cresol, thereby depositing precipitating a rubbery polymer. This rubbery polymer was

separated, washed with methanol and then dried in vacuo at the normal temperature. The thus obtained vinyl-cispolybutadiene rubber had a percent yield of 80 %. Thereafter, this vinyl-cis-polybutadiene rubber was treated with boiling n-hexane, and an insoluble matter and a soluble matter were separated and dried. The resulting boiling n-hexane soluble polymer had an ML of 31, a T-cp of 57 and a T-cp/ML relation of about 1.8 and had a microstructure consisting of 1.0 % of the 1,2-structure, 0.9 % of the trans-1,4-structure and 98.1 % of the cis-1,4structure. Furthermore, the [n] was 1.7. The number of monodispersed fiber crystals having a minor axis length of not more than 0.2 µm, which were contained in the vinvlcis-polybutadiene rubber, was 100 or more, the aspect ratio was not more than 10, and the melting point was 202 °C. [00921

(Production of vinyl-cis-polybutadiene sample 2)

A vinyl-cis-polybutadiene was obtained in the same manner as in the production process of the sample 1, except for not adding the unsaturated high-molecular substance (additive).

[0093]

Physical properties of the foregoing sample 1 and sample 2 are shown in Table 1.

[0094]				
Table 1				
	Sample Name		Sample 1	Sample 2
-High-	High-molecular substance		IR	Nil
(Am	(Amount of addition)		(10 % by weight)	weight)
Poly	Polymerization solvent		Cyclohexane	1
(SP value o	(SP value of polymerization solvent)	olvent)	8.1	\
	Mooney viscosity		31	+
4	[μ]		1.7	1.4
Characteristics of	Viscosity in toluene solution (cp)	ne solution (cp)	57	ţ
polybutadiene rubber		Cis	98.1	↓
100000000000000000000000000000000000000	Wicrostructure	Trans	6.0	+
	(0)	Vinyl	1.0	+
	Melting point of fiber crystal (°C)	iber crystal (°C)	202	+
	Number of mono	Number of monodispersed fiber		
	crys	crystals:		
	(Number of crysta	(Number of crystals having a minor	100 or more	4
the system to the contraction of 10-	axis length of no	axis length of not more than 0.2 µ		
polybutadiene	per 4	per 400 µ ²)		
crystalline fiber	Aspect ratio of monodispersed fiber	nodispersed fiber	Not more than	1
1	crystal		10	70 07 111014
	Major axis size of	Major axis size of crystalline fiber	0 0 +0 0 5	1 00000
	dispersed in matrix	×	0.2 00 0.3	T OT TIOTE
	Major axis size of	Major axis size of crystalline fiber	Not more than	
	dispersed in high-	dispersed in high-molecular substance	0.1	
% by weight of 1,2-polybutadiene	utadiene		12	↓
Number of monodispersed fiber crystals	fiber crystals		Large	Small

[0095]

[0096]

(Examples 1-1 to 1-4) and (Comparative Examples 1-1 to 1-3) By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 2 were kneaded by using a 1.7-L Banbury mixer for testing, thereby obtaining a kneaded material which is a silica compounded rubber composition for tire. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 $^{\circ}\mathrm{C}$. Next, this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 $^{\circ}\mathrm{C}$ for 30 minutes. The results are summarized and shown in Table 2.

The compositions of the Examples realize a small die swell, high abrasion resistance and low fuel consumption while keeping wet skid properties as compared with the case of using commercially available high cis-polybutadiene as in the Comparative Examples. On the other hand, in the compositions of the Comparative Examples, when the vinylcis-polybutadiene which does not meet the characteristics of the invention is used or the amount of addition of silica is low, although a small die swell and high abrasion resistance are realized, a remarkable lowering of the wet properties is caused so that the desired characteristics cannot be obtained. [0097]

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		Exan	Example		Сотра		Example
	1-1	1-2	1-3	1-4	1-1	1-2	1-3
S	Sample 1	Sample 1	Sample 1	Sample 1	1	Sample 2	Sample 1
	35	35	45	35	1	35	35
		25	15	1	1	-	-
	1	ı	1	1	35	1	1
	65	40	40	65	65	65	65
	27	27	27	21.5	27	27	9
	30	30	30	35	30	30	_
	9	9	9	7	9	9	1
Γ	15	15	15	1.5	15	15	15
-,	2	2	2	2	5	22	٣
2		2	2	2	2	2	2
1		1	1	1	1	1	1
1.3	3	1.3	1.3	1.4	1.3	1.3	1
0.7	7	7.0	7.0	6.0	7.0	0.7	ı
2		2	2	21	2	2	1.8
73		7.9	77	7.8	100	92	68
172	21	160	180	163	100	160	185
66		86	86	100	100	91	06
8	89	68	79	83	100	103	110
	ı						

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note 4) X50S (Equal-amount mixture of N330 and Si69, manufactured by Degussa) $\,$

Si69: Bis(3-triethoxysilylpropyl)-tetrasulfide (Note 5) Antioxidant: ANTAGE AS (a reaction product between amine and ketone)

(Note 6) Vulcanization accelerator: NOCCELER CZ (N-cyclohexyl-2-benzothiazole sulfenamide) (Note 7) Vulcanization accelerator: NOCCELER D (N,N'-diphenylquanidine)

[0098]

(Examples 2-1 to 2-5) and (Comparative Examples 2-1 to 2-4)
By using each of the foregoing sample 1 and sample 2,
compounding agents other than a vulcanization accelerator
and sulfur in a compounding preparation as shown in Table 3
were kneaded by using a 1.7-L Banbury mixer for testing,
thereby obtaining a kneaded material which is a rubber
composition for sidewall. On this occasion, a maximum
kneading temperature was adjusted at 170 to 180 °C. Next,
this kneaded material was kneaded with a vulcanization
accelerator and sulfur on a 10-inch roll, and the mixture
was rolled in a sheet form and then charged in a die for
vulcanization, thereby obtaining a vulcanizate. The
vulcanization was carried out at 150 °C for 30 minutes.
The results are summarized and shown in Table 3.

[00991 The compositions of the Examples are greatly improved with respect to the elastic modulus and highly balanced between the die swell and the exothermic characteristic as compared with the case of using commercially available high cis-polybutadiene as in the Comparative Examples. On the other hand, in the compositions of the Comparative Examples, when the commercially available BR is used, the die swell is large; and when the compounding amount of carbon black is high, the exothermic characteristic is deteriorated. Furthermore, when vinyl-cis-polybutadiene which is not satisfied in the invention is used or the amount of use of vinyl-cis-polybutadiene of the invention is low, the improvement effects did not reached the expected levels.

[0100]

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			Example			ŭ	Comparative Example	ve Examp	a
compounting capte	2-1	2-2	2-3	2-4	2-5	2-1	2-2	2-3	2-4
Kind of vinyl-cis-							,		,
poly-	Sampre	Sample	эашрте	Sample	Sample	1	Sample	Sample	Sample
butadiene	7	7	7	-	7		7	-	-
Amount (parts)	09	09	09	80	40	ı	09	09	10
NR (Note 1)	40	40	40	20	09	40	40	40	90
BR (Note 2)	1	-	-	1	-	09	1	-	i
Carbon black N330	40	55	30	40	40	40	40	7.0	40
Aromatic oil	10	10	10	10	10	10	10	10	10
Zinc oxide	т	3	3	3	3	m	e	e	6
Stearic acid	2	2	2	2	2	2	2	2	2
Antioxidant (Note 3)	1	1	1	1	1	П	1	П	1
Vulcanization	1.0	1.0	1.0	6.0	8.0	1.0	1.0	1.0	0.7
Sulfur	5	5		1 7	0 0	ر ب	r		2.2
Dhira i an I among at i an					2		2	2	7.7
of compounded									
material:									
Die swell index	73	64	84	89	83	100	06	28	86
Physical properties									
of vulcanizate:									
300 % tensile									
elastic modulus	190	240	160	165	185	100	135	269	104
(index)									
Exothermic									
characteristic (index)	80	06	69	75	98	100	104	107	101

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note 4) X50S (Equal-amount mixture of N330 and Si69, manufactured-by Degussa)

- Si69: Bis(3-triethoxysilylpropyl)-

tetrasulfide(Note 5) Antioxidant: ANTAGE AS (a reaction product between amine and ketone) (Note 64) Vulcanization accelerator: NOCCELER (N-

cyclohexyl-2-benzothiazole sulfenamide) -(Note 7) - Vulcanization - accelerator: NOCCELER - D (N, N'-

diphenylquanidine)

(Examples 3-1 to 3-4) and (Comparative Examples 3-1 to 3-5) By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 4 were kneaded by using a 1.7-L Banbury mixer for testing, thereby obtaining a kneaded material which is a rubber composition for cap tread of passenger automobile tire. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 °C. Next, this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 °C for 30 minutes. The results are summarized and shown in Table 4.

[0102]

The compositions of the Examples realize a high elastic modulus and high abrasion resistance while keeping wet skid properties as compared with the case of using commercially available high cis-polybutadiene as in the On the other hand, in the Comparative Examples. compositions of the Comparative Examples, when the vinylcis-polybutadiene which does not meet the characteristics of the invention is used or the amount of addition of SBR is low, although a high elastic modulus and high abrasion resistance are realized, a remarkable lowering of the wet properties is caused so that the desired characteristics cannot be obtained.

[0103]

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0 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2								,	
Compounding table		Exar	Example			Compar	Comparative Example	<pre>cample</pre>	
componing carry	3-1	3-2	3-3	3-4	3-1	3-2	3-3	3-4	3-5
Kind of vinyl-cis-	Sample	Sample	Sample	Sample		Sample		Sample	Sample
poly-	' ⊢	ı,	• ←	' ⊢	ı	5	ı	•	·
Amount (parts)	35	35	45	20	,	35	,	35	65
NR (Note 1)	1	25	15	40	35		ı	55	1
BR (Note 2)	1	-	ı	-	,	-	35	-	
SBR1500	65	40	40	40	65	65	65	10	35
Carbon black N330	09	09	09	09	09	09	09	09	09
Aromatic oil	15	15	15	15	15	15	15	15	15
Zinc oxide	3	٣	Э	m	e	m	e	m	m
Stearic acid	2	2	2	2	2	2	2	2	2
Antioxidant (Note 3)	1	1	1	П	П	П	1	н	1
Vulcanization	1.5	1.2	1.3	1.2	1.4	1.5	1.5	1.0	1.3
Sulfur	1.8	1.9	1.8	2.0	2.1	1.8	1.8	2.0	1.7
Physical properties of compounded									
material:	7.0	0	7.0		5	S	0	c	5
Physical properties	2	0	7/	0	TOO	25	TOT	0.6	70
of vulcanizate:									
300 % tensile									
elastic modulus (index)	183	174	190	151	100	126	88	160	215
Pico abrasion (index)	177	163	183	152	100	164	146	138	201
Wet skid properties (index)	66	16	96	86	100	91	88	94	93

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note-4) X50S (Equal-amount-mixture of N330 and Si69, manufactured by Degusa)

Si69: Bis(3-triethoxysilylpropyl)-tetrasulfide(Note 5)Antioxidant: ANTAGE AS (a reaction product between amine and ketone)

product between amine and ketone)
(Note 64) Vulcanization accelerator: NOCCELER CZ (Ncvclohex)-2-benzothiazole sulfenamidel-(Note 7)

cyclohexyl-2-benzothiazole sulfenamide) (Note 7) - Vulcanization accelerator: NOCCELER D (N.N'-diphenylquanidine)

[0104]

(Examples 4-1 to 4-3) and (Comparative Examples 4-1 to 4-2) By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 5 were kneaded by using a 1.7-1 Banbury mixer for testing, thereby obtaining a kneaded material which is a rubber composition for cap tread of large-sized vehicle tire. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 °C. Next, this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 °C for 30 minutes. The results are summarized and shown in Table 5.

[0105]

The compositions of the Examples realize a small die swell and excellent wet skild properties and abrasion resistance while keeping a high elastic modulus as compared with the case of using commercially available high cispolybutadiene as in the Comparative Examples. On the other hand, in the compositions of the Comparative Examples, since the vinyl-cis-polybutadiene which does not meet the characteristics of the invention is used, the abrasion resistance is not improved or a lowering of the wet skid properties is caused so that the desired characteristics cannot be obtained.

[0106]

Table 5

Table					
> det = paipainoumo		Example		Comparati	Comparative Example
componium cante	4-1	4-2	4-3	4-1	4-2
Kind of vinyl-cis-	Sample 1	Sample 1	Sample 1	1	Sammle 2
polybutadiene	1	1	1		
Amount (parts)	30	20	30	1	30
NR (Note 1)	7.0	80	7.0	7.0	7.0
BR (Note 2)	1	-	1	30	-
Carbon black N330	09	09	50	09	09
Aromatic oil	10	10	10	10	10
Zinc oxide	e	3	е	m	ю
Stearic acid	2	2	2	2	2
Antioxidant (Note 3)	1	1	П	П	1
Vulcanization accelerator	0 0	0	o c	c	c
(Note 4)	0.0	0.0	0.	0.0	°:
Sulfur	1	1	1	1	П
Physical properties of					
compounded material:					
Die swell index	74	82	80	100	95
Physical properties of					
vulcanizate:					
300 % tensile elastic	173	7 11 1		0	000
modulus (index)	C/T	# C T	177	T00	971
Pico abrasion (index)	163	144	125	100	133
Wet skid properties (index)	105	104	105	100	94

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note 4) X50S (Equal-amount mixture of N330 and Si69, manufactured by Degussa)

5:69: Bis(3-tricthoxysilylpropyl)-tetrasulfide(Note-5)Antioxidant: ANTAGE AS (a reaction product between amine and ketone)

(Note 64) Vulcanization accelerator: NOCCELER CZ (N-cyclohexyl-2-benzothiazole sulfenamide) (Note 7)

Vulcanization accelerator: NOCCELER D (NrN-diphenyl-yuanidine)

[0107]

(Examples 5-1 to 5-5) and (Comparative Examples 5-1 to $\underline{\mathtt{and}}$ 5-42)

By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 6 were kneaded by using a 1.7-L Banbury mixer for testing, thereby obtaining a kneaded material which is a high-hardness compounded rubber composition. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 °C. Next, this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 °C for 30 minutes. The results are summarized and shown in Table 6.

[0108]

The compositions of the Examples are greatly improved with respect to the elastic modulus and highly balanced between the die swell and the low exothermic characteristics. On the other hand, in the compositions of the Comparative Examples, when the commercially available BR is used, the die swell is large; and when the compounding amount of carbon black is high, the exothermic characteristic is deteriorated. Furthermore, when vinyl-cis-polybutadiene which is not satisfied in the invention is used or the amount of use of vinyl-cis-polybutadiene of the invention is low, the improvement effects did not reached the expected levels.

[0109]

Table 6

Compounding table			Example			Compa	Comparative Example
	5-1	5-2	2-3	5-4	5-5	5-1	5-2
Kind of vinyl-cis-	Camp lo 1	Sample	Sample	Sample	1 olamo		Sample
polybutadiene	Sampre 1	Н	1	1	T andmed	ı	\vdash
Amount (parts)	50	7.0	30	50	50	1	20
NR (Note 1)	50	30	7.0	50	50	50	50
BR (Note 2)	1	1	1	-	-	50	ı
Carbon black N330	7.0	7.0	7.0	09	08	7.0	7.0
Aromatic oil	5	5	5	10	10	22	2
Zinc oxide	D.	5	2	e	3	5	2
Stearic acid	2	2	2	2	2	2	2
Antioxidant (Note 3)	1	1	1	1	Τ	1	1
Vulcanization accelerator	1.0	1.2	0.8	1.0	1.0	1.0	1.0
(Note 4)							
Sulfur	3.0	2.8	3.2	3.0	3.0	3.0	3.0
Physical properties of							
compounded material:							
Die swell index	7.5	68	85	83	65	100	68
Physical properties of							
vulcanizate							
Hardness (index)	124	134	16	116	130	100	109
Low exothermic	c	C D	C	c c		00	0
characteristics (index)	00	٠,	6	00	# O	700	# O T

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note 4) X50S (Equal-amount mixture of N330 and Si69, manufactured by Degussa)

Si69: Bis(3-triethexysilylpropyl)-tetraculfide(Note-5)Antioxidant: ANTAGE AS (a reaction product between amine and ketone)

[0110]

(Examples 6-1 to 5-5) and (Comparative Examples 6-1 to 6-5) By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 7 were kneaded by using a 1.7-L Banbury mixer for testing, thereby obtaining a kneaded material which is a high-hardness compounded rubber composition. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 °C. Next, this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 °C for 30 minutes.

The compositions of the Examples are greatly improved with respect to the elastic modulus and highly balanced between the die swell and the exothermic characteristic. On the other hand, in the compositions of the Comparative Examples, when the commercially available BR is used, the die swell is large; and when the compounding amount of carbon black is high, the exothermic characteristic is deteriorated. Furthermore, when vinyl-cis-polybutadiene which is not satisfied in the invention is used or the amount of use of vinyl-cis-polybutadiene of the invention is low, the improvement effects did not reached the

The results are summarized and shown in Table 7.

[0112]

expected levels.

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Compounding table		Ιſ	Example			,	Comparative		Example	
	T-9	7-9	6-3	5-9	6-5	1-9	7-9	6-3	6-4	6-5
	Sam-	Sam-	Sam-	Sam-	Sam-		Sam-	Sam-	Sam-	Sam-
	ple 1	ple 1	ple 1	ple 1	ple 1		ple 2	ple 1	ple 1	ple 1
	50	7.0	30	50	20	1	20	20	20	10
	50	30	7.0	50	20	20	20	20	20	90
	1	1	1	1	1	50	1	1	1	1
	40	40	40	50	35	40	40	09	20	40
	10	10	10	10	10	10	10	10	10	10
	2	2	2	2	2	2	S	2	2	5
	2	2	2	2	2	2	2	2	2	2
	1	1	1	1	П	7	1	-	,	1
	1.0	0.9	0.8	1.0	1.0	1.0	1.0	1.0	1.0	7.0
	1.5	1.7	2.0	1.5	1.5	1.5	1.5	1.5	1.5	2.2
	71	63	85	62	78	100	88	55	103	94
	210	234	160	233	190	100	138	260	113	120
	83	79	88	96	75	100	901	110	87	66

(Note 1) NR: RSS#1 (Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.) (Note 3) Ultrasil VN3 GR (manufactured by Degussa) (Note 4) X50S (Equal-amount mixture of N330 and Si69, manufactured-by-Degussa) Si69: Bis(3-triethoxysilvlpropyl)-

tetrasulfide (Note 5) Antioxidant: ANTAGE AS (a reaction product between amine and ketone) (Note 64) Vulcanization accelerator: NOCCELER CZ cvclohexvl-2-benzothiazole sulfenamide) (Note 7) - Vulcanization accelerator: NOCCELER D (N,N'diphenylguanidine)

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(Examples 7-1 to 7-3) and (Comparative Examples 7-1 to 7-2) By using each of the foregoing sample 1 and sample 2, compounding agents other than a vulcanization accelerator and sulfur in a compounding preparation as shown in Table 8 were kneaded by using a 1.7-L Banbury mixer for testing, thereby obtaining a kneaded material which is a rubber composition for tire cord coating. On this occasion, a maximum kneading temperature was adjusted at 170 to 180 °C. this kneaded material was kneaded with a vulcanization accelerator and sulfur on a 10-inch roll, and the mixture was rolled in a sheet form and then charged in a die for vulcanization, thereby obtaining a vulcanizate. The vulcanization was carried out at 150 °C for 30 minutes. The results are summarized and shown in Table 8.

[0114] The compositions of the Examples have a small die swell, are largely improved with respect to the green modulus while keeping a high elastic modulus and are excellent in adhesiveness to metals. On the other hand, in the compositions of the Comparative Examples, since the vinvl-cis-polybutadiene which does not meet characteristics of the invention is used or commercially available high cis-polybutadiene is used, the desired characteristics cannot be obtained. [0115]

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Commonation table		Example		Comparativ	Comparative Example
compounding capie	7-1	7-2	7-3	7-1	7-2
Kind of vinyl-cis- polybutadiene	Sample 1	Sample 1	Sample 1	-	Sample 2
Amount (parts)	35	20	35	1	35
NR (Note 1)	65	80	65	65	65
BR (Note 2)	1	ı	1	35	1
Carbon black N330	09	09	50	09	09
Zinc oxide	7	7	7	7	7
Stearic acid	2	2	2	2	2
Cobalt stearate	3	3	m	3	8
Antioxidant (Note 3)	2	2	2	2	2
Vulcanization accelerator (Note 4)	0.8	0.8	8.0	0.8	0.8
Sulfur	1.5	1.5	1.5	1.5	1.5
Physical properties of					
compounded material:	C	,	· ·		
Die swell (index)	27	181	80	100	96
100 % green modulus (index)	136	T74	113	TOO	TOS
Physical properties of					
vulcanizate:					
300 % tensile elastic	17.4	6	120	001	106
modulus (index)	T / T	100	027	TOO	TC 0
Tensile strength (index)	163	144	125	100	138
Adhesive strength to metal	107	106	105	100	86
(lndex)					

(Note 2) BR: Polybutadiene (UBEPOL-BR150, manufacture by Ube Industries, Ltd.)

(Note 3) Ultrasil VN3 GR (manufactured by Degussa)

(Note 4) X50S (Equal-amount mixture of N330 and Si69_r manufactured by Degussa)

Si69.

Bis(3-triethoxysilylprepyl)tetraculfide(Note-5)Antioxidant: ANTAGE AS (a reaction
product between amine and ketone)

Vulcanization accelerator: NOCCELER D (N,N'-diphenylguanidine)

Industrial Applicability

[0116]

The silica compounded rubber composition for tire which is used in the invention has a small die swell and excellent extrusion processability and moldability while keeping high wet skid properties, improves the workability of tire manufacture and has excellent abrasion resistance and low fuel consumption.

The rubber composition for sidewall of the invention has low fuel consumption with respective to a vulcanizate thereof and has a small die swell.

The rubber composition for passenger automobile tire according to the invention is able to realize a high elastic modulus and high abrasion resistance while keeping wet skid properties.

The rubber composition for large-sized vehicle tire according to the invention has a small die swell and excellent extrusion processability while keeping a high elastic modulus and is able to be highly balanced between wet skid properties and abrasion resistance.

The high-hardness compounded rubber composition according to the invention improves dimensional stability at the time of processing and durability of tire at the same time while keeping a high hardness and is able to make the both performances compatible with each in a good balance.

The rubber composition for base tread according to the invention improves die swell and low fuel consumption at the same time and is able to make the both performances compatible with each in a good balance.

The rubber composition for tire cord coating according to the invention has a small die swell, large green strength and excellent extrusion processability and moldability while keeping a high elastic modulus, has excellent adhesiveness to metals and is able to be highly

balanced among the respective characteristics.